DRAWINGS

5'	GTG	GGC	ATC				. 610	- (1)	3 C10	GIG	CIG	GIG	ATC	.GCG	CGG	GTG	CGC	54 CGG
	<u>Val</u>	Gly	Met	: Val	Gly	Asn	Va]	Leu	<u>.</u> Leu	ı Val	Leu	Val	Ile	Ala	Arm	 Val	7~~	Arg
	CIG	CAC	63 AAC	GIG	ACG	72 AAC	TTC	CIC	81 ATC	GGC	AAC	CIG 30	GCC	TTG	99 TCC	GAC	GIG	108 CTC
	Leu	His	Asn	Val	Thr	Asn	Phe	Leu	Ile	Gly	Asn	Leu	Ala	Leu	Ser	Aso	 Val	 T.e.i
	ATG	TGC	117 ACC	GCC	TGC	126 GTG	CCC	CTC	135 ACG	CTG	GCC	144 TAT	GCC	TTC	153 GAG	CCA	CGC	162 GGC
	Met	Cys	Thr	Ala	Cys	Val	Pro	Leu	Thr	Leu	Ala	Tyr	Ala	Phe	Glu	Pro	Ara	 Glv
	TGG	GIG	171 TTC	GGC	GGC	180 GGC	CIG	TGC	189 CAC	CIG	GTC	198 TTC	TTC	.: CTG	207 CAG	CCG	GTC	216 ACC
	Trp	Val	Phe	Gly	Gly	Gly	Leu	Cys	His	Leu	Val	Phe	Phe	Leu	 Gln	Pro	 Val	 Thr
	GIC	TAT	225 GTG	TCG	GIG	234 TTC	ACG	CTC	243 ACC	ACC	ATC	252 GCA	GIG	GAC	261 CGG	TAC	GTC	270 GTG
		-3-	vai	Ser						Thr	Ile	Ala	Val	Asp .	Arg	Tyr	Val	Val
(CIG	GIG	CAC	CCG	CTG	288 AGG	CGG	CGC	297 ATC	3'								1
j	Leu	Val	His	Pro	Leu .	Arg	Arg	Arg	Ile									

that the first of the first of the first of the state of the first of

18 27 36 GGC CTG CTG GTC ACC TAC CTG CTC CCT CTG GTC ATC CTG TCT TAC Gly Leu Leu Val Thr Tyr Leu Leu Pro Leu Leu Val Ile Leu Leu Ser Tyr 63 72 81 90 GTC CGG GTG TCA GTG AAG CTC CGC AAC CGC GTG GTG CCG GGC TGC GTG ACC CAG --- --- --- --- --- --- --- --- --- --- --- --- --- ---Val Arg Val Ser Val Lys Leu Arg Asn Arg Val Val Pro Gly Cys Val Thr Gln 117 126 135 144 AGC CAG GCC GAC TGG GAC CGC GCT CGG CGC CGC ACC TTC TGC TTG CTG GTG --- --- --- --- --- --- --- --- --- --- --- --- --- ---Ser Gln Ala Asp Trp Asp Arg Ala Arg Arg Arg Thr Phe Cys Leu Leu Val 180 189 GTG GTC GTG GTG TTT GCC ATC TGC TGG TTG CCT TAC TAC 3' Val Val Val Val Phe Ala Ile Cys Trp Leu Pro Tyr Tyr

Fig. 3

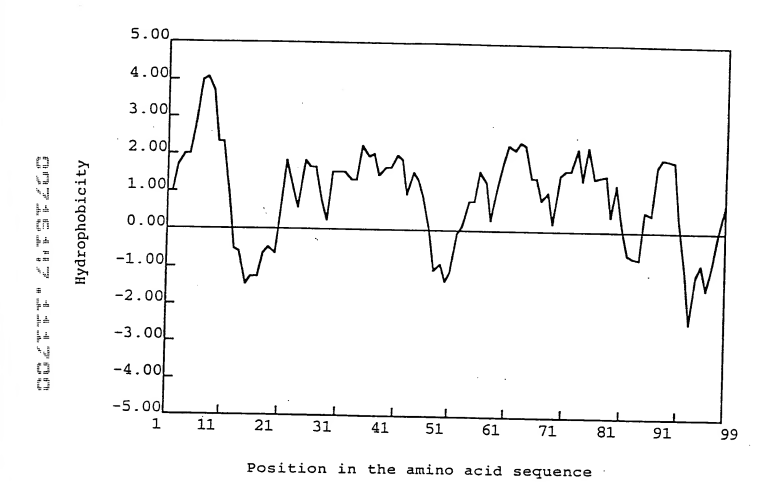
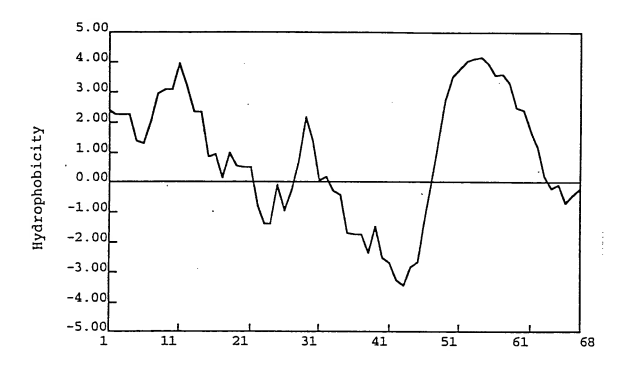


Fig. 4

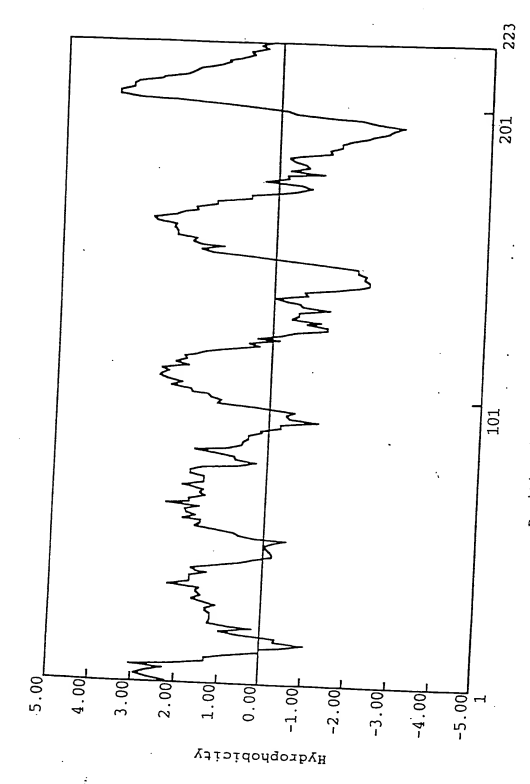


Position in the amino acid sequence

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10 20 30 40 1 VCMVCNVLLV LVITARVRRLH NVTNFELTONL ALSDVLMCTA CVPLTLAYAF 1 LGVSGNLALI IIILKQKEMR NVTNFELTONL SFSDLLVAVM CLPFIFVYIL	60 70 80 90° 100 EPRGWVFG3G LCHLVFFFLQP WIVYWSVFTL TTIAVDRYVV LVHPLRRRI- MDH-WWFG5T MCKLNPFVQC WSITWSIFSL VLIAVERHQL IINPRGWRPN	140	IWVLAVASSL PFVIYQILID EPFQNVSLAA FKDKYVCFDK	160 170 180 190 200GLDLV TYILPLEVIL LSY VRVSVKERNR VVPGCVTQSQ FPSDSHRESY TILLVEQYF GPLCFIFICY FKIYIREKRR NNWMDKIRDS	240
30 NVTNELICAL NVTNELICAL	80 VIVYVSVETE VSITVSIFSE	130	PFVIYQILTD	180 LSV GPLCFIFICY	230 VFALCWLPYY -FAVCWIPLT
20 LV TARVRRLH II TUKQKEMR	70 LCHINFFLOP MCKINPFVQC	120	IWVLAVASSL	170 TYTELEVIL	220 TFCLLAVWW NVMLESIWA
10 Venvenvlev Levsenlali	60 EPRG <mark>WVFG</mark> 3G MDH-MVFGET	110	101 NRHAYIGITV	160 GLUUV FPSDSHRUSY	210 220 230 201 ALWDRARRIRY TECLLAVIVIV VEALOWIPPY 201 KYRSSETKRI NVMLLSIVVA -FAVOWIPLT
ਜਜ	51	101	101	151	201 201
p19P2 S12863	p19P2 S12863	p19P2	<u>\$12863</u>	p19P2 S12863	p19P2 S12863

5' GTG GGC ATG GTG GGC AAC ATC CTG GTG GTG ATC GCG GGG GTG CGC CGG
Val Gly Met Val Gly Asn Ile Leu Leu Val Leu Val Ile Ala Arg Val Arg Arg
63
CTG TAC AAC GTG ACG AAT TTC CTC ATC GGC AAC CTG GCC TTG TCC GAC GTG CTC Leu Tyr Asn Val Thr Asn Phe Leu Ile Gly Asn Leu Ala Leu Ser Asp Val Leu
ATG TGC ACC GCC TGC GTG CCG CTC ACG CTG GCC TAT GCC TTC GAG CCA CGC GGC Met Cys Thr Ala Cys Val Pro Leu Thr Leu Ala Tyr Ala Phe Glu Pro Arg Gly
TGG GTG TTC GGC GGC GGC CTG TGC CAC CTG GTC TTC CTG CAG GCG GTC ACC
Trp Val Phe Gly Gly Leu Cys His Leu Val Phe Phe Leu Gln Ala Val Thr
225 234 243 252 261 270 GTC TAT GTG TCG GTG TTC ACG CTC ACC ACC ATC GCA GTG GAC CGC TAC GTC GTG Val Tyr Val Sor Val
Val Tyr Val Ser Val Phe Thr Leu Thr Thr Ile Ala Val Asp Arg Tyr Val Val
CTG GTG CAC CCG CTG AGG CGG CGC ATC TCG CTG CGC CTC AGC GCC TAC GCT GTG Leu Val His Pro Lou Air
Leu Val His Pro Leu Arg Arg Arg Ile Ser Leu Arg Leu Ser Ala Tyr Ala Val
333 342 351 360 362
CTG GCC ATC TGG GTG CTG TCC GCG GTG CTG GCG CTG CCC GCC G
TAT CAC GIT GAG CTT 336 405 414 423
TAT CAC GTG GAG CTC AAG CCG CAC GAC GTG CGC CTC TGC GAG GAG TTC TGG GGC TYT His Val Glu Leu Lys Bro His Annual Company Compan
Tyr His Val Glu Leu Lys Pro His Asp Val Arg Leu Cys Glu Glu Phe Trp Gly
TCC CAG GAG CGC CAG CGC CAG CTC TAC GCC TGG GGG CTG CTG GTC ACC TAC Ser Gln Glu Arg Gln Arg Gln Leu Tyr Ala Trp Gly Leu Leu Val Thr Tyr
495 504 513 522 The Tyr
CTG CTC CTG CTG GTC ATC CTC CTG TCT TAC GCC CGG GTG TCA GTG AAG CTC Leu Leu Pro Leu Leu Val Tla
549 CS2 Leu Leu Ser Tyr Ala Arg Val Ser Val Lys Leu
COC AAC CGC GTG GTG CCG GGC CGC GTG ACC CAG ACC GTG GTG S85
The Gly Arg Val Thr Gln Ser Gln Ala Asp Trp Asp Arg
CCT CCG CCC CCG CCC ACC TTC TCC TTC CTC GTC GTC GTC GTC GTC G
Ala Arg Arg Arg Thr Phe Cys Leu Leu Val Val Val Val Val Val Phe Thr
657 666
Leu Cys Trp Leu Pro Phe Phe
sed FIO FUE PRE

50	100	150	200	250 250 250
50 CVPLTLAYAF CVPLTLAYAF	100 LVHPLRRRI LVHPLRRRIS	150 EFWGSQERQR	200 QSQADWDRAR QSQADWDRAR	250
40 ALSDVLWCTA ALSDVLWCTA	90 TTIAVDRYVV TTIAVDRYVV	140 LKPHDVRLCE	190 RNRVVPGCVT RNRVVPGRVT	240
30 NVTNFLIGNL NVTNFLIGNL	80 VTVYVSVFTL VTVYVSVFTL	130 PAAVHTYHVE	180 LSYVRVSVKL LSYARVSVKL	230 PFF
20 LVIARVRRDH LVIARVRRDY	70 LCHINFFLQP LCHLVFFLQA	120 IWVLSAVLAL	170 TYLLPLLVIL TYLLPLLVIL	220 VVVVEALCML VVVVEIT.CWL
10 VGMVGNVILLV VGMVGNILLLV	60 EPRGWVFGGG EPRGWVFGGG	110 LRLSAYAVLA	160 QLYAWGILILIV	RRRTFCLLWV
H H	51	101	151	201
 p19P2 pG3-2/pG1-10	p19P2 pG3-2/ p G1-10	p19P2 pG3-2/pG1-10	p19P2 pG3-2/pG1-10	p19P2 pG3-2/pG1-10



Position in the amino acid sequence

1 CATCGTCAAGCAGATGAAGATCATCCACGAGGATGGCTACTCCGAGGGCCAGCAGAAATT	F 60
61 CTGCCCCTTCTTCCCGCGAGTGCTTTCCCGCTCTCCAAACCCCACTCCCAGGTGGCCATG	1
121 GCCTCATCGACCACTCGGGGCCCCAGGGTTTCTGACTTATTTTCTGGGCTGCCGCCGGCG 1 AlaSerSerThrThrArgGlyProArgValSerAspLeuPheSerGlyLeuProProAla	21
181 GTCACAACTCCCGCCAACCAGAGCGCAGAGGCCTCGGCGGGCAACGGGTCGGTGGCTGGC	240 41
241 GCGGACGCTCCAGCCGTCACGCCCTTCCAGAGCCTGCAGCTGGTGCATCAGCTGAAGGGG 41 AlaAspAlaProAlaValThrProPheGlnSerLeuGlnLeuValHisGlnLeuLysGly	300 61
301 CTGATCGTGCTGCTCTACAGCGTCGTGGTGGTCGTGGGGGCAACTGCCTGC	360 81
361 GTGCTGGTGATCGCGGGGTGCGCCGGCTGCACAACGTGACGAACTTCCTCATCGGCAAC 81 ValLeuValIleAlaArgValArgArgLeuHisAsnValThrAsnPheLeuIleGlyAsn	420 101
421 CTGGCCTTGTCCGACGTGCTCATGTGCACCGCCTGCGTGCCGCTCACGCTGGCCTATGCC 101 LeuAlaLeuSerAspValLeuMetCysThrAlaCysValProLeuThrLeuAlaTyrAla	480 121
481 TTCGAGCCACGCGGCTGGGTGTTCGGCGGCGCCTGTGCCACCTGGTCTTCTTCCTGCAG 121 PheGluProArgGlyTrpValPheGlyGlyGlyLeuCysHisLeuValPhePheLeuGln	540 141
541 CCGGTCACCGTCTATGTGTCGGTGTTCACGCTCACCACCATCGCAGTGCACCGCTACGTC 141 ProvalThrValTyrValSerValPheThrLeuThrThrIleAlaValAspArgTyrVal	600 161
501 GTGCTGGTGCACCCGCTCAGGCGCGCATCTCGCTGCGCCTCAGCGCCTACGCTGCTGCTG 161 ValLeuValHisProLeuArgArgIleSerLeuArgLeuSerAlaTyrAlaValLeu	660 181
561 GCCATCTGGGCGCTGTCCGGGGTGCTGGCGCTGCCGCCGTGCACACCTATCACGTG 181 AlaIleTrpAlaLeuSerAlaValLeuAlaLeuProAlaAlaValHisThrTyrHisVal	720 201
721 GAGCTCAAGCCGCACGACGTGCGCCTCTGCGAGGAGTTCTGGGGGCTCCCAGGAGCGCCAG 201 GluLeuLysProHisAspValArgLeuCysGluGluPheTrpGlySerGlnGluArgGln	780 221
781 CGCCAGCTCTACGCCTGGGGGGCTGCTGCTCACCTACCTGCTCCCTCTGCTCGTCATC 221 ArgGlnLeuTyrAlaTrpGlyLeuLeuLeuValThrTyrLeuLeuProLeuLeuVallle	840 241
841 CTCCTGTCTTACGTCCGGGTGTCAGTGAAGCTCCGCAACCGCGTGGTGCCGGGCTGCGTG 241 LeuLeuSerTyrValArgValSerValLysLeuArgAsnArgValValProGlyCysVal	900 261
: 901 ACCCAGAGCCAGGCCGACTGGGACCGCGCGCGCGCGCGCG	960 281
961 GTGGTCGTGGTGTTCGCCGTCTGCTGCTGCCGCTGCACGTCTTCAACCTGCTGCGG 281 ValValValValValPheAlaValCysTrpLeuProLeuHisValPheAsnLeuLeuArg	1020 301
1021 GACCTCGACCCCACGCCATCGACCCTTACGCCTTTGGGCTGGTGCAGCTGCTCTGCCAC 301 AspLeuAspProHisAlaIleAspProTytAlaPheGlyLeuValGlnLeuLeuCysHis	1080 321
1081 TGGCTCGCCATGAGTTCGGCCTGCTACAACCCCTTCATCTACGCCTGGCTGCACGACAGC 321 TrpLeuAlaMetSerSerAlaCysTyrAsnProPheIleTyrAlaTrpLeuHisAspSer	1140 341
1141 TTCCGCGAGGAGCTGCGCAAACTGTTGGTCGCTTGGCCCCGCAAGATAGCCCCCCATGGC 341 PheArgGluGluLeuArgLysLeuLeuValAlaTrpProArgLysIleAlaProHisGly	1200 361
1.201 CAGAATATGACCGTCAGCGTGGTCATCTGATGCCACTTAGCCAGGCCTTGGTCAAGGAGC 361 GlnAsnMetThrValSerValValIle***	1260
1261 TCCACTTCAACTGGCCTCCTAGGGCACCACTCGAGGTCAATCTGGTGCTTATTCTCAGCA	371 1320
1321 CCAGAGCTAGC 371	371 1331
	371

Fig. 10

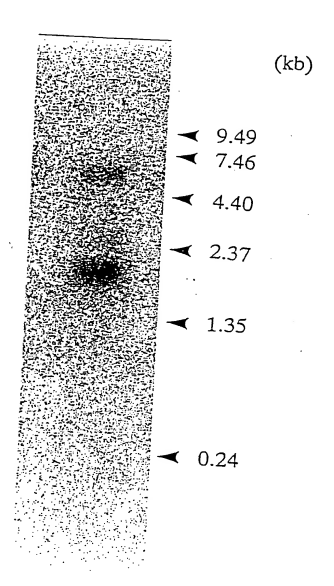
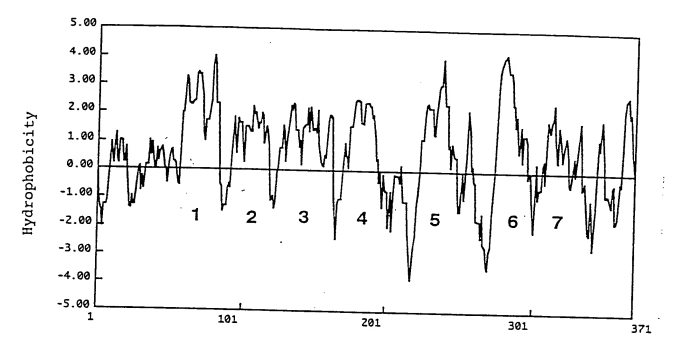


Fig. 11



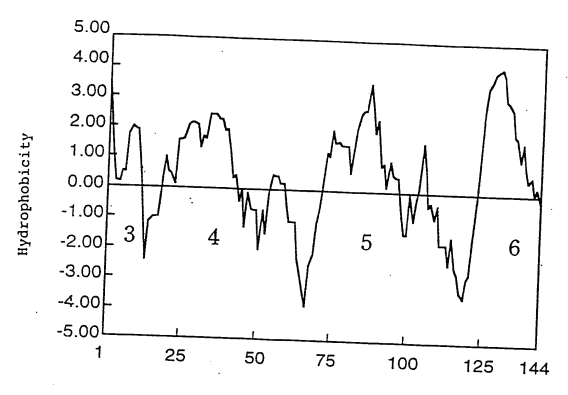
Position in the amino acid sequence

5'	CTG	TG	T (9 STC	ATC	GC	:	.8 .G G2		~~	4.00	C G1		2 T.T.	CIC	GI	G CA	c c c	JG (7)	'A CC	54 T CGG
	reu	CV	<u> </u>	<u>'ZI</u>	<u></u>	A L	a Va	l As	D A	rg	Ty	r Va	1 V	al	Leu	ı Va	l Hi	s Pr	o Le	u Ar	r CGG g Arg
	CGC	AT	r 1	CA A	CTG	AGO	7 CT	2 C AG	C G	CC	81 TAC	I CGC	GG	TG	90 CTG	GGC	: ATY	9 Э	9	т ст	108
	Arg	Ile	∍ S	er	Leu	Arg	Le	u Se	r A	la	Tyr	Al	a V	al	Leu	Gly	 ' Ile	 • Tr	 D Al.	 a To	A TCT u Ser
	GCA	GTC		TG	GCG	CTG	12	s GCC	C G	CG	135 GTG	CA(C Ac	CC	144 TAC	CAT	' GTG	15	3 3	~ aa	162
	ALG	vai	. L:	eu	Ala	Leu	Pro	o Ala	a Ai	la	Val	His	s Ti	ır	Tyr	His	Val	Gli	ı Let	Lys	Pro
	CAC	GAC	G:	IG	AGC	CTC	180 TGC) CGAC	GP	4G	189 TTC	TGC	GC	€C	198 TCG	CAG	GAG	207	7 - Car	· ~~	216
:	HIS	Asp	Vā	il	Ser	Ļeu	Cys	Glu	G1	u.	Phe	Tre	Gl	y .	Ser	Gln	Glu	Arc	Gln	Aro	Gln
	ATC	TAC	GC	:5 :C '	TGG	GGG	234 CTG	CTI	CI	ਹ ਹ	243 GGC	ACC	TA	T :	252 ITG	CTC	ccc	261 CTG		CCC	270
	+15	17.7	<u> </u>	٠ .	ırp	GIY	Leu	Leu	Le	u (Gly	Thr	Ty:	r I	Leu	Leu	Pro	Leu	Leu	Ala	Ile
1	CTC	CIG	TC	9 T 1	EAC	GTA	288 CGG	GTG	TC	A (297 FIG	AAG	CI	3 G A	106 166 .	AAC	CGC	315 GTG	GTG	ررس	324 GCC
1	Leu !	Leu	Se	r 1	ŊΥ,	Val	Arg	Val	Sez	- V	/al	Lys	Leu	ı A	rg 1	Asn	Arg	Val	 Val	Pro	Gly
2	AGC (TG	33. AC	3 C C	AG 2	AGT	342 CAA	GCT	GAC	3 T	51 GG (GAC	CGA	3 4 G	60 CG (CT (CGC	369 CGC	CCC	۸ حت	378
S	Ser (/al	Th	G	ln s	Ser (Gln	Ala	Asp	T	tb 7	Asp	Arg	- A	 la <i>1</i>	ira i	 Ara	 Aro	 2rc		 Dh -
1	CT C	TG	CIC	G	TG G	TG (396 GTG	GTG	GTA	40 G	05 IG 7	ITC	ACG	4: C:	14 IC 1	GC 1	rgg (423	CCC	ייייי	432
	ys L	eu	Leu	. Va	al V	al v	/al	Val	Val	Va	al <u>E</u>	Phe	Thr	Le	eu C	ys 1	ן מֹז.	Leu	Pro	 Phe	 Tyr

CT 3'

50 50 -30	100 100 21	150 150 71	200 200 200 121	250 250 171
SO CVPLTILAYAF CVPLTILAYAF	100 LVHPLRRRI LVHPLRRRIS LVHPLRRRIS	150 EFWGSQERQR EFWGSQERQR	200 QSQADWDRAR QSQADWDRAR QSQADWDRAR	250
40 L ALSDVLMCTA L ALSDVLMCTA	90 TTIAVDRYVV TTIAVDRYVV CVIAVDRYVV	140 LKPHDVRLCE LKPHDVSLCE	190 RNRVVPGCVT RNRVVPGRVT RNRVVPGSVT	240
30 H NVTANFLIGNI	80 VTVYVSVFTL VTVYVSVFTL	130 PAAVHTYHVE PAAVHTYHVE	180 LSYVRVSVKL LSYARVSVKL LSYVRVSVKL	230 Phr Phr
20 V INTARVRRDH V LVIARVRRDY	TOHENFELOP	120 IWALSAVLAL IWALSAVLAL	170 TYLLPLLVIL TYLLPLLVIL TYLLPLLAIL	220 WWWFTLCWL WWWFTLCWL
VGMVGNT	60 EPRGMVFGGG EPRGMVFGGG	110 LRLSAYAVLA LRLSAYAVLS	160 QLYAWGLLLV QLYAWGLLLV QLYAWGLLLG	RRRTFCLLVV RRRTFCLLVV RRRTFCLLVV
1 1 - 79	51 51 -29	101 101 22	151 151 72	201 201 122
p19P2 pG3-2/pG1-10 p5S38	p19P2 pG3-2/pG1-10 p5S38	p19P2 pG3-2/pG1-10 p5S38	p19P2 pG3-2/pG1-10 p5S38	p19P2 pG3-2/pG1-10 p5s38

Fig. 14



Position in the amino acid sequence

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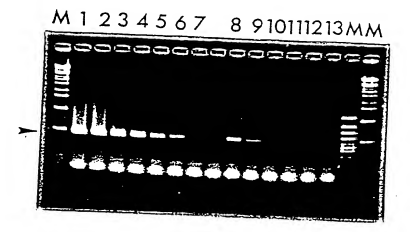
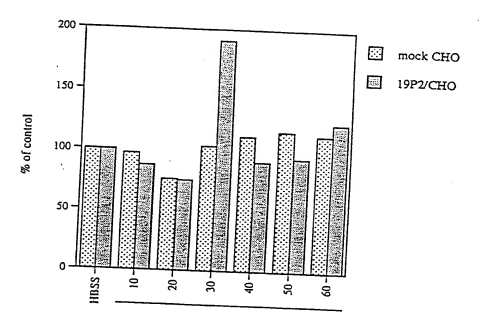
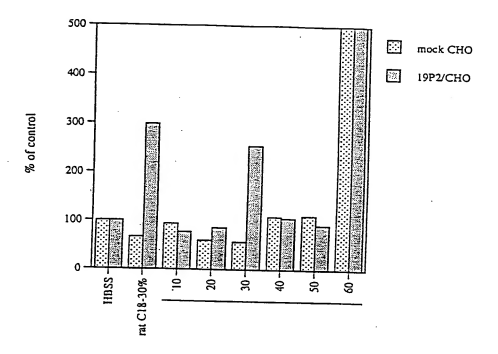


Fig. 16

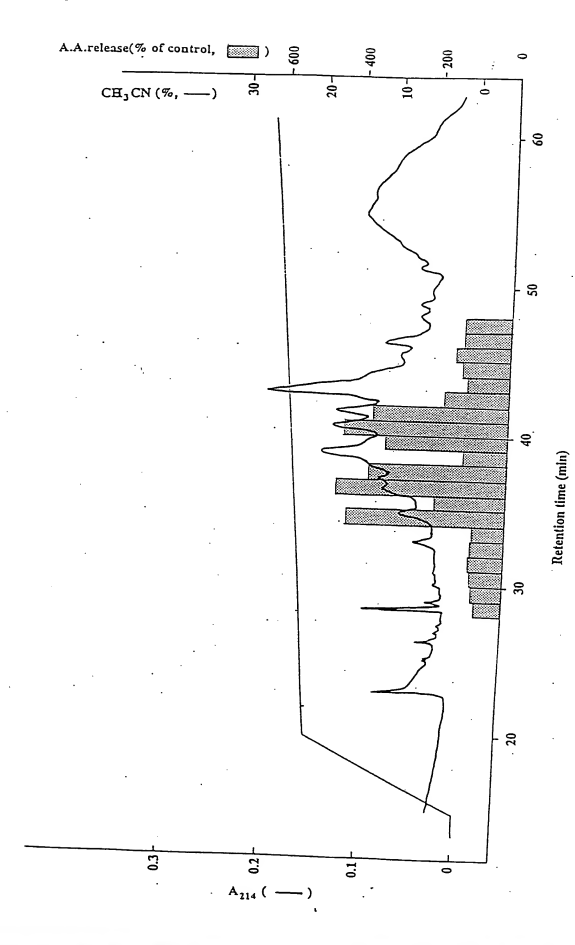


rat whole brain extract C_{18} -column CH_3CN elution (%)

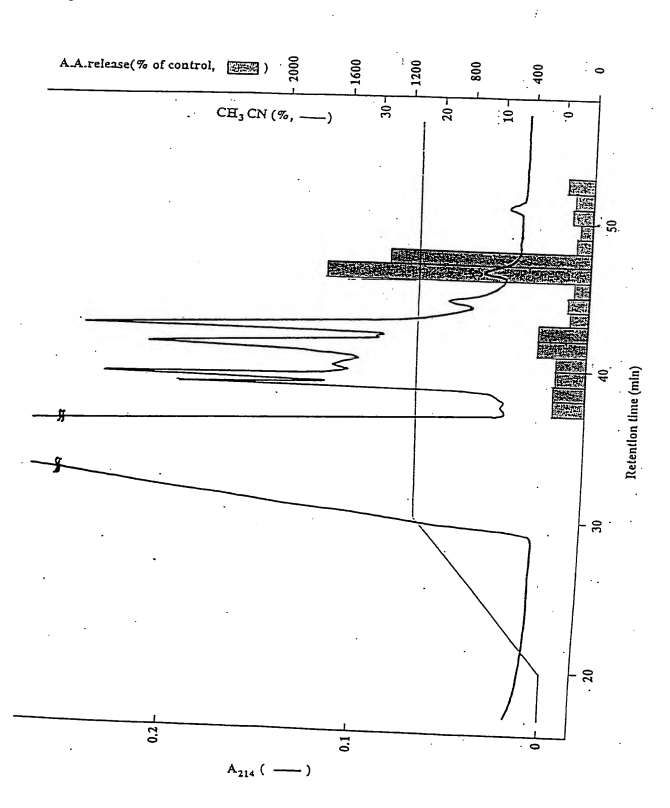
Fig. 17

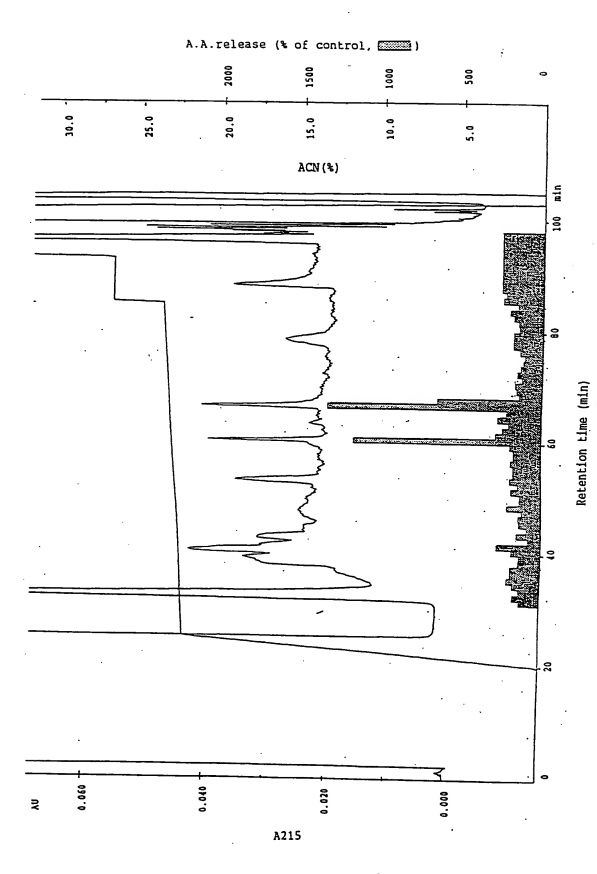


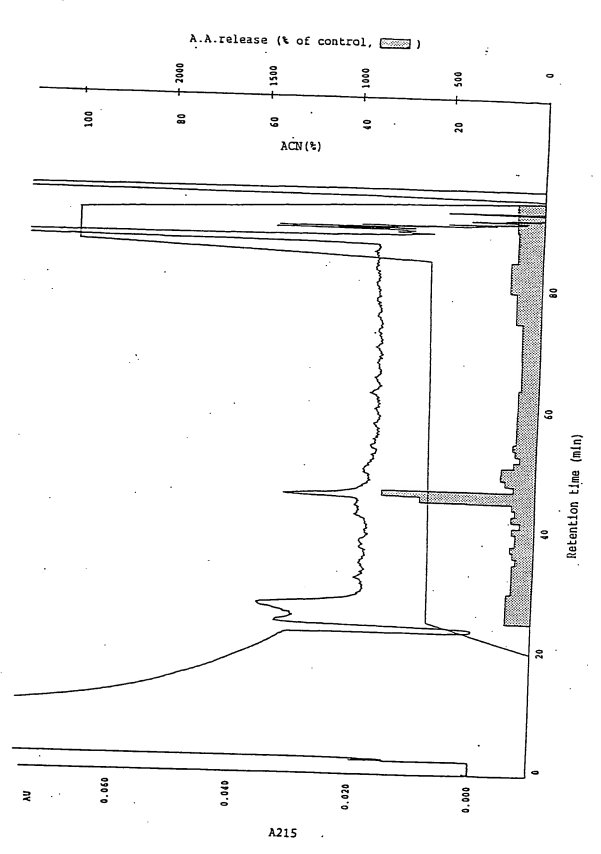
bovine hypothalamus extract C_{18} -column CH_3CN elution (%)











) – T					_											
			9			18			27			36			45			54
5 '	GCC	CAC	CAG	CAC	TCC	ATG	GAG	ATC	CGC	ACC	CCC	GAC	ATC	AAC	CCT	GCC	TGG	TAC
	Ala	His	Gln	His	Ser	Met	Glu	Ile	Arg	Thr	Pro	Asp	Ile	Asn	Pro	Ala	Trp	Tyr
•																	-	•
			63			72												
	GCG	GGC	CGT	GGG	ATC	CGG	CCC	G 3	•									
								-										
	Ala	Gly	Arg	Gly	Ile	Arg	Pro											
		3	P3-2	?														

1	GTGGAATGAAGGCGGTGGGGCCTGGCTCCTCTGCCTGCTGCTGCTGGCCCTG MetLysAlaValGlyAlaTrpLeuLeuCysLeuLeuLeuLeuGlyLeuAlaLeu	59 18
60 19	CAGGGGGCTGCCAGCAGAGCCCACCAGCACTCCATGGAGATCCGCACCCCCGACATCAAC GlnGlyAlaAlaSerArgAlaHisGlnHisSerMetGluIleArgThrProAsplleAsn	119 38
	CCTGCCT	
39	ProAla	126
		40

L GIGGAATGAAGGCGGTGGGGCCTGGCTCCTCTGCCTGCTGCTGCTGGGCCTGGCCTGGCCTG	
2 delighed LeuCysteuLeuLeuGlyteuAlateu	5: 1:
CAGGGGGCTGCCAGCAGCCCAGCAGC	-
GlnGlyAlaAlaSerArgAlaHisGlnHisSerMetGluIleArgThrProAspIleAsp	119
CCTGCCTGGTACGCPCCCCCTGGG	3.8
ProAlaTrpTyrAlaGlyArgGlyIleArgProValGlyArgDroCo	179
<u> </u>	58
GCCCCGGGGGACGCAGGCCTGGCCCCGGCGTGTCCCGGGGGGGG	
AlaProGlyAspGlyProArgProGlyProArgArgValProAlaC	239
andydifioAlaCysPheArgLeuGlu	78
GGCGGYGCTGAGCCCTCCCGAGCCCTCCCGGGGCGCTGACCCCGG	
GlyGlyAlaGluProSerArgAlaLeuProGlyArgLeuThraleGl	299
The control of the co	98
TAACAGCGGGAGCCTGCCCCACCCCTCCTCCACCACCACCACCACCACCACCACCA	
TO THE OWN OF THE OWN OF THE OWN OF THE OWN OWN OF THE OWN	359
33/11333	98
AATAAAAGCAGCTGGCTTGTT	
	380
· ·	98
	MethysalavalglyalatrpheuleuCysLeuLeuLeuGlyLeuAlaLeu CAGGGGGCTGCCAGCAGCACCACCAGCACTCCATGGAGATCCGCACCCCGACATCAAC GlnGlyAlaAlaSerArgAlaHisGlnHisSerMetGlulleArgThrProAspIleAsn CCTGCCTGGTACGCRGGCCGTGGGATCCGGCCGGCGGCGGAAGAGCT ProAlaTrpTyrAlaGlyArgGlyIleArgProValGlyArgPheGlyArgArgArgAla GCCCCGGGGGACCCAGGCCTGGCCCCGGGGGCTGCTTCCGCCTGGAA AlaProGlyAspGlyProArgProGlyProArgArgValProAlaCysPheArgLeuGlu GGCGGYGCTGAGCCCTCCCGAGCCTCCCGGGGCGGCCAGCTGGTCCAGGAA GlyGlyAlaGluProSerArgAlaLeuProGlyArgLeuThrAlaGlnLeuValGlnGlu TAACAGCGGGAGCCTGCCCCCCACCCCTCCTCCACCAGCCACCTTCCCTCCAGTCCT

Fig. 24(b)

1	GTGGAATGAAGGCGGTGGGGCCTGGCTCCTCTGCCTGCTGCTGGGCCTGGCCCTG	
- 1	MetLysAlaValGlyAlaTrpLeuLeuCysLeuLeuLeuLeuGlyLeuAlaLeu	59
- 0		18
00	CAGGGGGCTGCCAGCAGCACCACCACCATGGAGATCCGCACCCCGACATCAAC	
19	GlnGlyAlaAlaSerArgAlaHisGlnHisSerMetGluIleArgThrProAspIleAsn	119 38
120	TOTOGOTACOCKOGCCGTGGGATCCCGCCCCCCCCCCCCCCCCCCCCCCCCC	
39	ProAlaTrpTyrAlaGlyArgGlyIleArgProValGlyArgPheGlyArgArgArgAla	179
		58
180		
59	AlaLeuGlyAspGlyProArgProGlyProArgArgValProAlaCysPheArgLeuGlu	239
	·,	78
240	GGCGGYGCTGAGCCCTCCCGAGCCCTCCCGGGGCGGCTGACGGCCCAGCTGGTCCAGGAA	
79	GlyGlyAlaGluProSerArgAlaLeuProGlyArgLeuThrAlaGlnLeuValGlnGlu	299
		98
300	TAACAGCGGGAGCCTGCCCCCCCCCCCCCCCCCCCCCCC	
98	The state of the s	359
		98
	AATAAAAGCAGCTGGCTTGTT	300
98		380
		98

Fig. 25

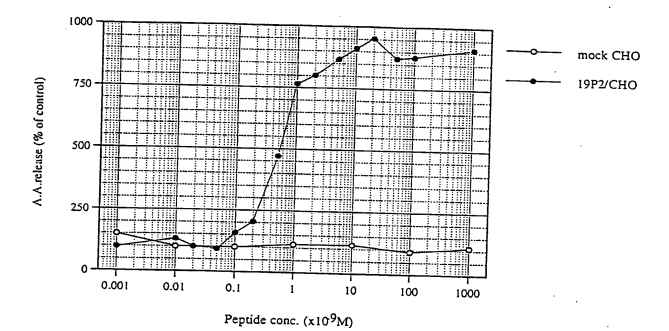


Fig. 26

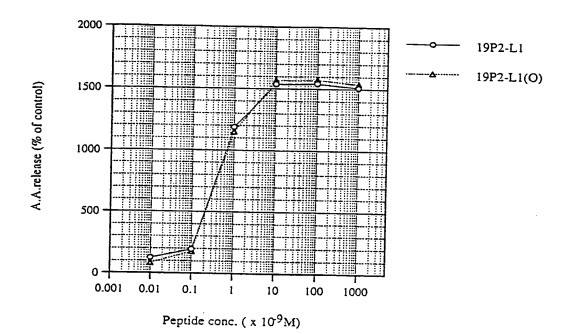
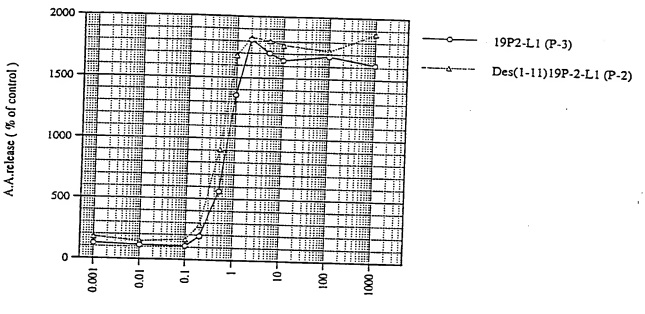


Fig. 27



Peptide conc. (x 109M)

Fig. 28



Fig. 29

10	20	30	40	50	60
ATGAAGGCGG	TGGGGGCCTG	GCTCCTCTGC	CIGCIGCIGC	TGGGCCTGGC	60 CCTGCAGGGG
70	80	90	100	110	120
GCIGCCAGCA	. GHOCCCHCCH	. GCACTCCA1G	GAGATCCGCA	GTGAGTGTCT	AGCCCCGCCC
130	140	150	160	170	180
	0001414466	6666666	CCACTICCIG	GGCIGGGACA	TCCTTGCTAA
190	200	210	220	230	240
GCAICCIGGG	GTTGGGGTTT				
250	260	270	280	290	300
	Graddiccc	GGCCCAGCAC	ACGGGGGAGG	GTCACTCCTC	ACCACACGGG
310	320	330	340	350	360
1000010000	CTGAGTGCAC				
370	380	390	400	410	420
	CCTGGTGTGA				
430	440	450	460	470	480
	CGGGTGAACC				
490	500	510	520	530	540
	AGCTGAGCAC				
550	560	570	580	590	600
	GCGCTCTTCT				
AGGCCGTGGG	620 ATCCGGCCCG	630	640	650	660
ACCCAGGCCT	680	690 CTCTTCCCCC	700	710	720
	GCCCCCGGC				
	740	750	760	770	780
	CTCCCGGGGC	COCTONCOCC	CCAGCIGGTC	CAGGAATAA.	• • • • • • • • • •

Fig. 30

	· · · · · · · · · · · · · · · · · · ·	
genome cDNA	10 20 30 40 50 1 ATGAAGGGG TGGGGGCCTG GCTCCTCTGC CTGCTGCTGC TGGGCCTGGC 1 ATGAAGGCGG TGGGGCCTG GCTCCTCTGC CTGCTGCTGC TGGGCCTGGC	50 50
genome cDNA	1 COLOCTORATE GETTER CHECK CHECK CCY CCYLL CYCLAS CHECKLES	100
genome cDNA	101	.50 .50
genome cDNA	29	00
genome cDNA	ZV1 ====================================	50 50
genome cDNA	ADI ====================================	00 00
genome CDNA	310 320 330 340 350 301 TGGCCTGGGG CTGAGTGCAC GTCACCCATG AGAACGGGGC TGTGAGGACA 35	
genome CDNA	360 370 380 390 400 351 GGAAAGGAAG GGGAGTGTGT CCTGGTGTGA GTCTGAAATC CTACTTCCCA 40 351	00
genome CDNA	410 420 430 440 450 401 AAGCCACCCC AGCACCAGAA ATGGGCGCTC CGGGTGAACC TCCTGTGCGG 45 401	50
genome CDNA	460 470 480 490 500 451 GTGGGTGGTC CTGGCCATGGC CTGGGCGACA GGCAGCCATG AGCTGAGCAC 50 451 50	
genome cDNA	501 550	-
genome cDNA	560 570 580 590 600 551 GCGCTCTTCT CTCTCTTCC AGCCCCCGAC ATCAACCCTG CCTGGTACGC 600 551	
genome cDNA	610 620 630 640 650 601 AGGCCGTGGG ATCCGGCCGG TGGGCCGCTT CGGCCGGGA AGAGCTGCCC 650 601 GGGCCGTGGG ATCCGGCCCG TGGGCCGCTT CGGCCGGCGA AGAGCTGCCC 650	
genome cDNA	660 670 680 690 700 651 TGGGGGACGG ACCCAGGCCT GGCCCCCGGC GTGTGCCGGC CTGCTTCCGC 700 651 CGGGGGACGG ACCCAGGCCT GGCCCCCGGC GTGTGCCGGC CTGCTTCCGC 700	
genome CDNA	710 720 730 740 750 701 CTGGAAGGCG GTGCTGAGCC CTCCCGAGCC CTCCCGGGGC GGCTGACGGC 750 701 CTGGAAGGCG GCGCTGAGCC CTCCCGAGCC CTCCCGGGGC GGCTGACGGC 750	
genome cDNA	760 770 780 790 800 751 CCAGCTGGTC CAGGAATAA. 800 751 CCAGCTGGTC CAGGAATAA. 800	

5'	ATG			GIG		GCC	TGG		27 CTC	TGC	CIG	36 CTG	CIG	CIG	45 GGC	CIG	GCC	54 : CTG
	PI	Λ.	A	V	G	A	W	L	L	С	L	L	L	L	G	L	A	L
	CAG	GGG	63 GCT	GCC	AGC	72 AGA	GCC	CAC	81 CAG	CAC	TCC	90 ATG	GAG	ATC	250 99	ACC	ccc	108 GAC
	Q	G	A	A	S	R	A	H	Q	H	s	M	E		R	т	 P	
	ATC	AAC	117 CCT	GCC	TGG	126 TAC	GCA	GGC	135 CGT	GGG	ATC	144 CGG	ccc	GTG	153 GGC	CGC	TTC	162 GGC
	I		P	A		Y.			R		I	R	P	v	G	 R	 F	·
	CGG	CGA	171 AGA	GCT	GCC	180 CTG	GGG	GAC	189 GGA	ccc	AGG	198 CCT	GGC	CCC	207 CGG	CGT	GTG	_
	R	R	R	A	A	L	G	D	G	P	R	P	G	p	 R	 R		 P
	GCC	TGC	225 TTC		CIG	234 GAA	GGC	GGT 	243 GCT	GAG	CCC	252 TCC	CGA	GCC	261 CTC	CCG	GGG	270 CGG
	A	С	F	R	L	E	G	G	Α	E	P	s	R	 A	L	P		 R
		ACG	GCC		CTG	288 GTC	CAG	GAA	297 TAA	3'								
	L	T,	A	Q	L	V	Q	E	*									

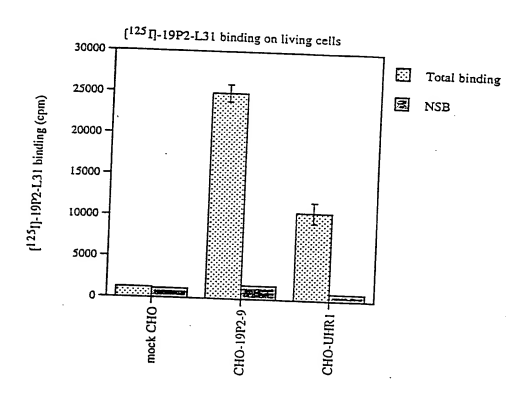
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1	Metal all out combined to the rectification of the	59
	MetAlaLeuLysThrTrpLeuLeuCysLeuLeuLeu	12
60	TIMEST TOO TOO TOO CAGGGGCTTCCAGCCGAGCCCACCAGCACTCCATCCACACAACA	119
13	LeuSerLeuValLeuProGlyAlaSerSerArgAlaHisGlnHisSerMetGluThrArg	32
120	THE CONTROL OF THE CO	179
33	ThrProAspIleAsnProAlaTrpTyrThrGlyArgGlyIleArgProValGlyArgPhe	52
180	TO THE STATE OF THE PROPERTY O	239
53	GlyArgArgAlaThrProArgAspValThrGlyLeuGlyGlnLeuSerCysLeuPro	
		72
240	CTGGATGGACGCACCAAGTTCTCTCAGCGTGGATAACACCCCAGCTCGAGAAGACAGTGC	299
73	LeuAspGlyArgThrLysPheSerGlnArgGly***	
		83
300	TGCTGAGCCCAAGCCCACACTCCCTGTCCCCTGCAGACCCTCCTCTACCCTCCCT	359
83		83
		0.5
360	CTGCT	364
83		02

bovine.aa	M 77 3 3 3 3	
bovine.seq	M K A V G A W L L	
rat.seq	-18	
rac.seq	1 GGCATCATCC AGGAAGACGG AGCATGG CCCTGAAGAC GTGGCTTCTG	32 50
bovine.aa	C L L L G L A L Q G A A S R A H	30
bovine.seq	70 00	
rat.seq		82
_	51 TECTTECTEC TECTAAGCTT GETCCTCCCA GEGGCTTCCA GCCGAGCCCA	100
bovine.aa	OHENDO	100
	INPAWYA	
bovine.seq		
rat.seq	83 CCAGCACTCC ATGGAGATCC GCACCCCCGA CATCAACCCT GCCTGGTACG	132
-	101 CCAGCACTCC ATGGAGACAA GAACCCCTGA TATCAATCCT GCCTGGTACA	150
bovine.aa	,	130
	162 GRRRAA	
bovine.seq	100 170 100	
rat.seg	133 CGGGCCGTGG GATCCGGCCC GTGGGCCGGCT TCGGCCGGCG AAGAGCTGCC	182
	TOGGCAGGAG ALGGCCAAGA	200
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	PGDG PRP GPR RVPA CFR	
bovine.seq	210 220 220	
rat.seq	183 CCGGGGGACG GACCCAGGCC TGGCCCCCGG CGTGTGCCGG CCTGCTTCCG	232
	201 CCGAGGGATG TCACTGGACT TGGCCAACTCA GCTGCCTCCC	250
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bovine.aa		300
2011116.22	Q L V Q E *	
bovine.seq	310 320 330 340 350	
	283 CCCAGCTGGT CCAGGAATAA CACCCCCAGG	•
rat.seq	The circles caree career and a second	332
	======================================	350
harrin -	360 370 380 390 400	
bovine.seq	333 TCCACCAGCC ACCTTCCCTC CAGTCCTAAT AAAACCAGCT COO	
rat.seq	Transcribed Ciccletter Terger	382
	***************************************	400

1	GGCCTCCTCGGAGGAGCCAAGGGATGAAGGTGCTGAGGGCCTGGCTCCTGTGCCTGCTG	
1	Maria de l'Addege Clege l'Ege	59
	MetLysValLeuArgAlaTrpLeuLeuCysLeuLeu	.12
60	ATGCTGGGCCTGCGGGGGGGGGGGCTGCAAGTCGTACCCATCGGCACTCCATGGAGATC	
13	MetLeuGlyLeuAlaLeuArgGlyAlaAlaSerArgThrHisArgHisSerMetGluIle	119
	1	32
120	COCACCCCIGACATCAATCCTGCCTGCTACGCCAGTCCCCCCATCACCCCCCCATCACCCCCCCC	
33	ArgThrProAspIleAsnProAlaTrpTyTAlaSerArgGlyIleArgProValGlyArg	179
	' ' ' ' ' ' ' ' ' ' ' ' ' ' ' ' ' ' '	52
180	TTCGGTCGGAGGGGGACCCTGGGGGACGTCCCCAAGCCTGGCCTGCGACCCCGGCTG	
53	PheGlyArgArgAlaThrLeuGlyAspValProLysProGlyLeuArgProArgLeu	239
	1 - 3 - 3 - 3 - 12 - 11 - 11 - 12 - 12 -	72
240	ACCTGCTTCCCCCTGGAAGGCGGTGCTATGTCGTCCCAGGATGGCTGACAGCCAGC	
73	ThrCysPheProLeuGluGlyGlyAlaMetSerSerGlnAspGly***	299
	1 ************************************	87
300	CAAGAAACTCACTCTGGAGCCTCCCCACCCCACCCTCTCCTCTCCTTCGGGCTCCTTTC	
87	The state of the s	359
		87
360	cc	
87		361
		87

bovine.aa rat.aa human.aa	10 20 30 40 50 1 MKAVGAWLLC LLLIGLAIQG AASRAHQHSM EIRTPDINPA WYAGRGIRPV 1 M-ALKIWLLC LLLLISLVIPG ASSRAHQHSM ETRTPDINPA WYTGRGIRPV 1 MKVLRAWILC LIMIGIAIRG AASRTHRHSM EIRTPDINPA WYASRGIRPV	50 50 50
bovine.aa rat.aa human.aa	60 70 80 90 100 51 GREGRRAAP GOGEREGERR VPACERLEGG AEPSRALPGR LTAQLVQE*. 51 GREGRRATE REVIGLG QUSCLPIDGR TRESORG*. 51 GREGRRATL GOVEREGERP RUTCEPLEGG AMSSQDG*.	100 100 100

Fig. 36



cells; 0.5 x 10⁷ cells/ml

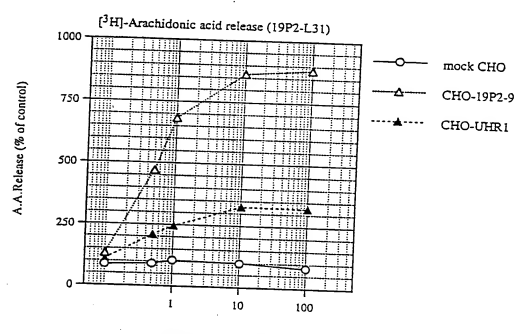
[125]]-19P2-L31; 200pM(avg.63857.3cpm) NSB; 200nM(x 1,000)

reaction; RT, 2.5hr

in HBSS + 0.05% BSA + 0.05% CHAPS

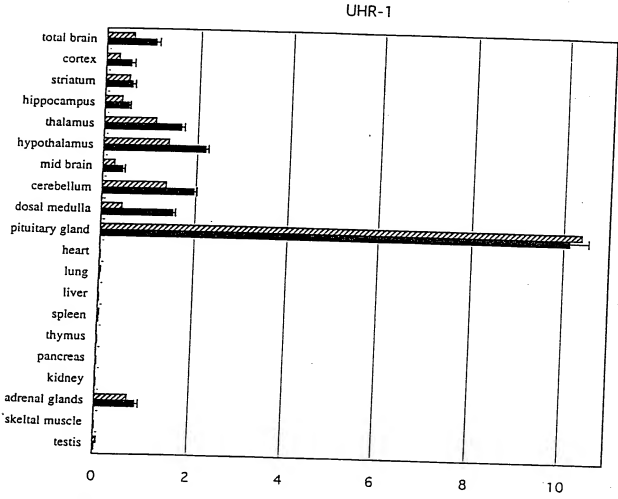
in 100 µl

Fig. 37



19P2-L31 (x 10-9M)

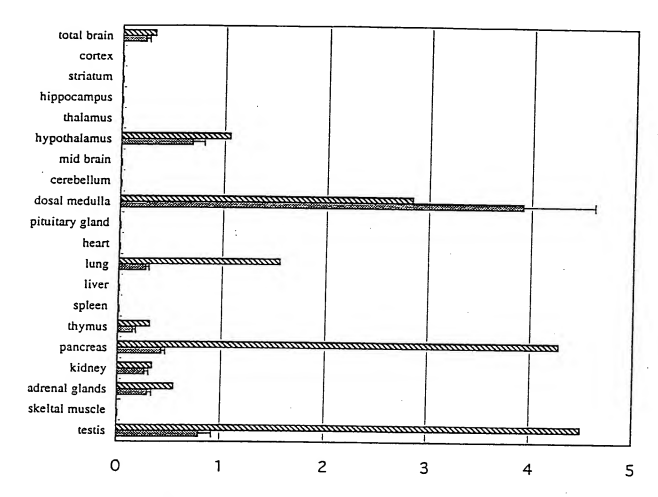
Fig. 38



■ receptor±S.E.M.(pgx10-3/4ng polyA+RNA) ② receptor/G3PDH(x2.5x10-2)

Fig. 39

ligand polypeptide



■ ligand \pm S.E.M.(pgx10-3/4ng polyA+RNA) \boxtimes ligand/G3PDH(x10-4)

Fig. 40

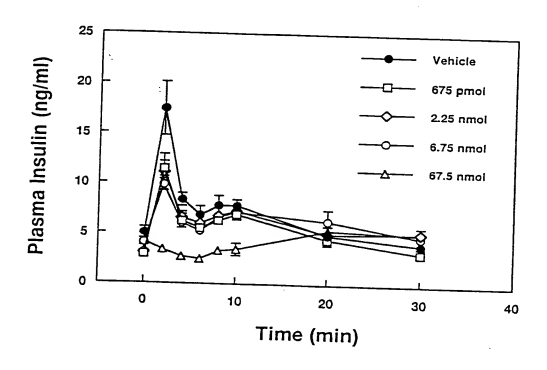
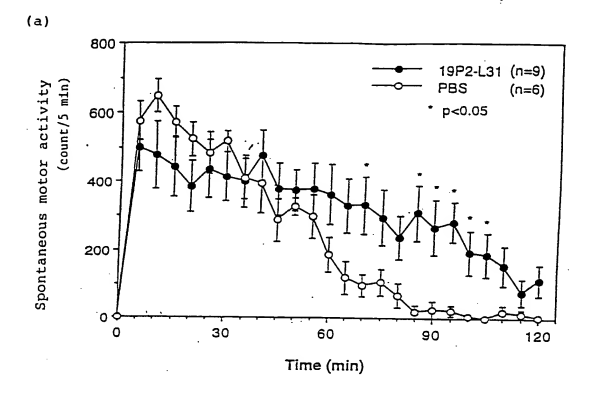


Fig. 41



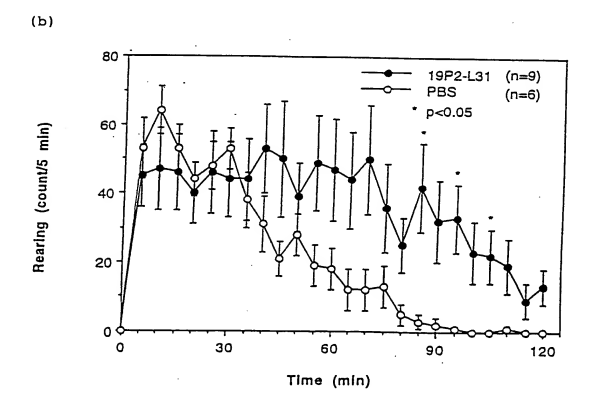
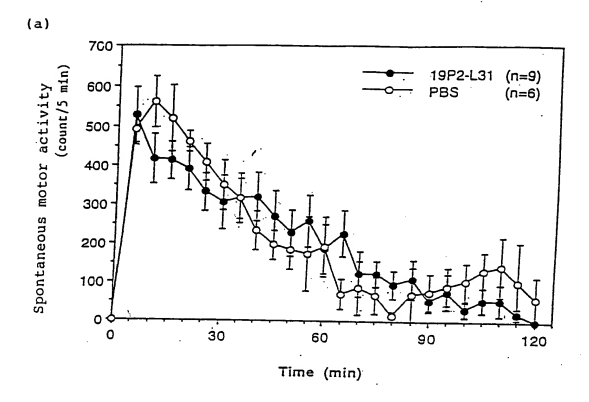


Fig. 42



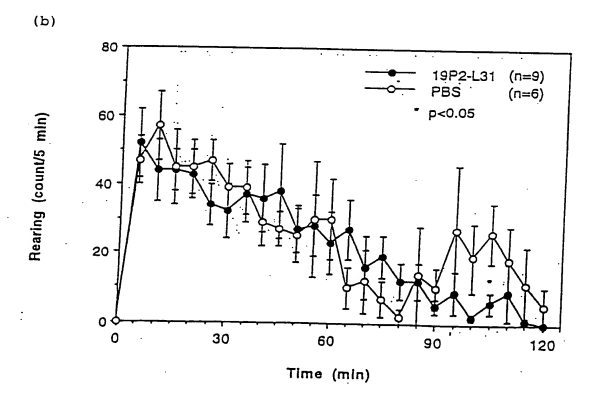
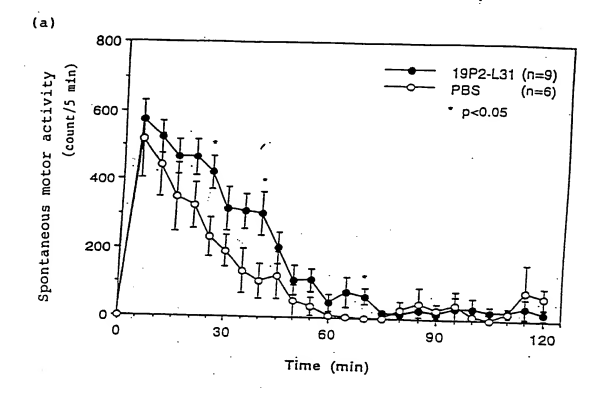


Fig. 43



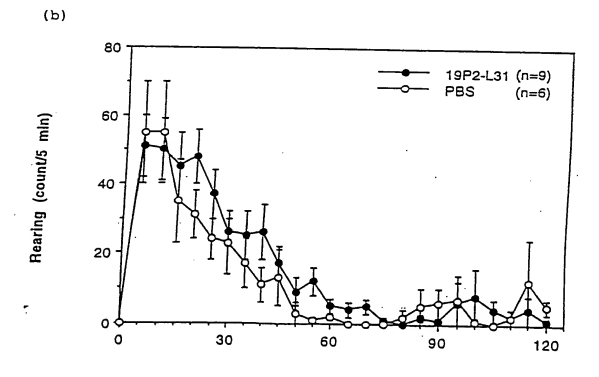
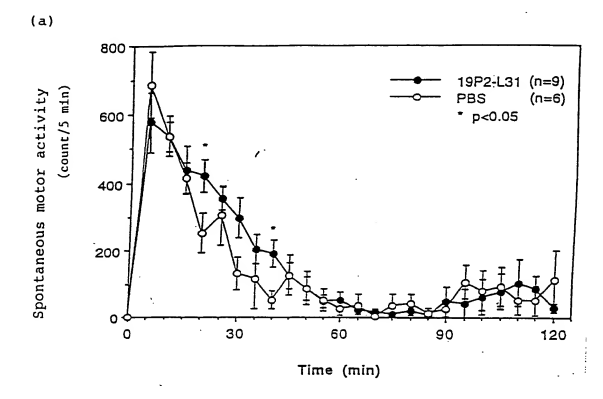


Fig. 44



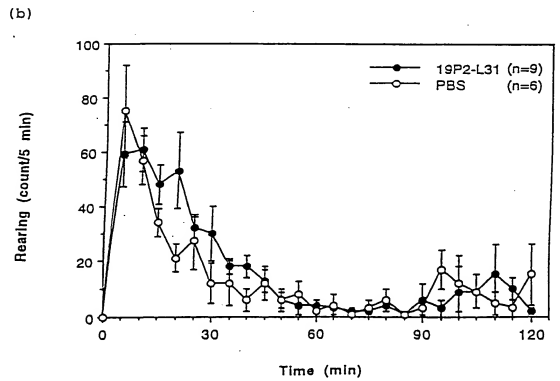
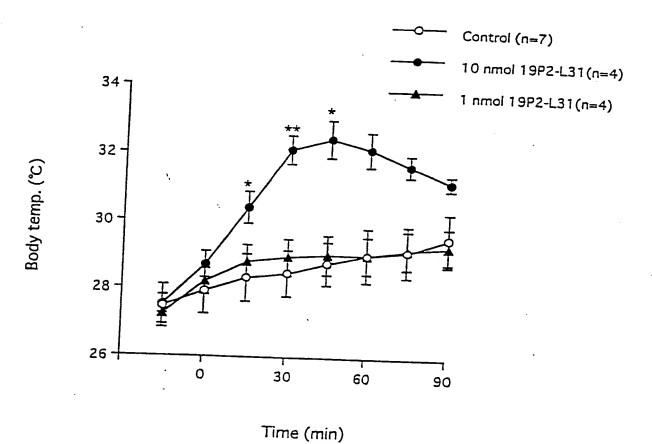
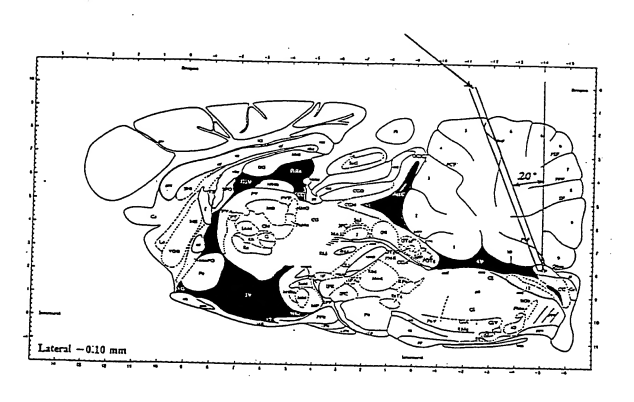


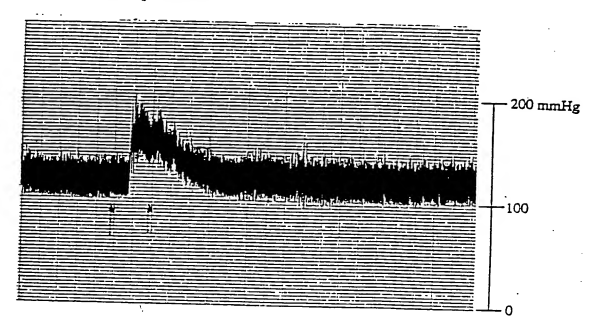
Fig. 45



micro-injection cannula



direct blood pressure



mean blood pressure

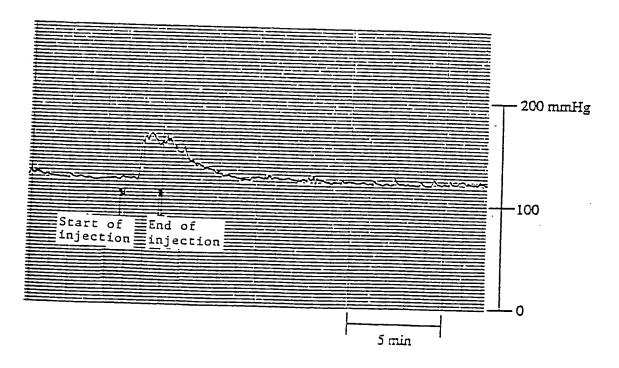


Fig. 48

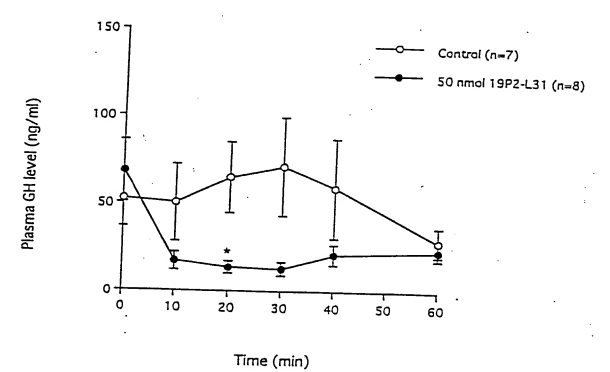
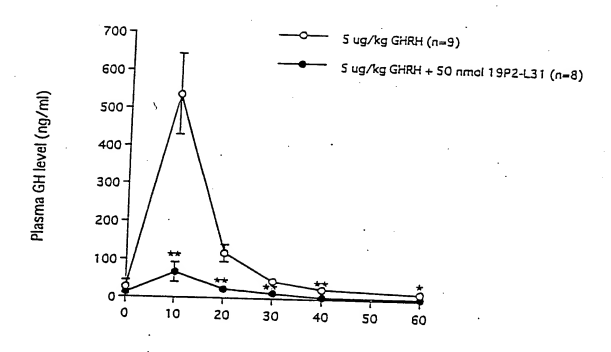
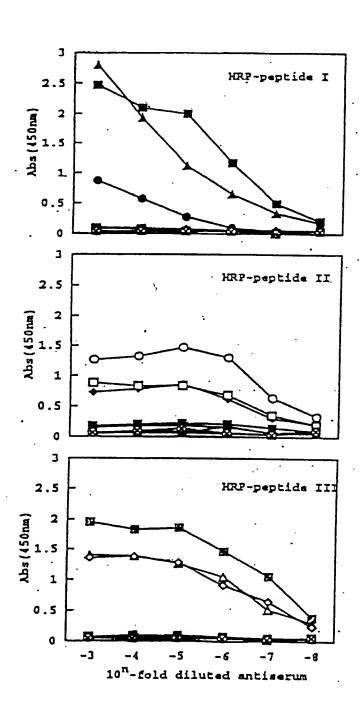


Fig. 49



Time (min)

Fig. 50



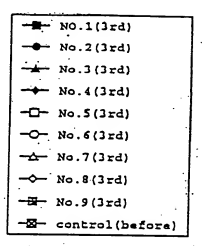
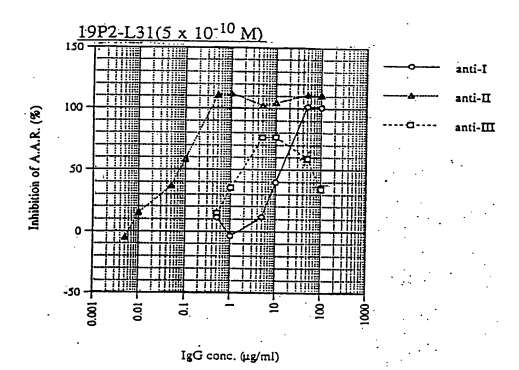


Fig. 51





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Leu Val His Cln Leu Lys Cly Leu Ile Val Mec Leu Tyc Ser Ile Val Val Val 333 342 351 360 3169 378 ARE THE COL COL TOT GIVE CIT CIT AGE CIT CIT AGE CIT CIT COL Het Cys Ala Ala Cys Val Pro Leu Thr Leu Ala Tyr Ala Phe Glu Pro Arg Gly 549 ... 558 ... 567 ... 576 ... 585 ... 594 ... 594 ... 500 .. ACT ACC TOC COC TOC DC AAC COC TTC ACC DAT COC TOC CAC CAC ACC TTC
Not See See Ale Cyn Tyr Am Pro Phe Ele Tyr Ale TTD Cou Mis Any See Phe OUT OND MATE AND AND OTHE AND OTH AND THE TRA TRA 1GITY Olin Assay Neet The Val See Val Val Ile *** ***

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